Appl. No. 09/635,864 Attorney Docket No.: 600-1-087CIP1CON

Page 28

EXHIBIT 3

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Blast 2 Sequences results

PubMed

Entrez

BLAST

OMIM

Taxonomy

Structure

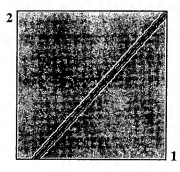
BLAST 2 SEQUENCES RESULTS VERSION BLASTP 2.2.15 [Oct-15-2006]

Matrix BLOSUM62 🔻	gap open: 11 gap extension: 1	a comment			
x_dropoff: 50 expect:	10.0000 wordsize: 3 Filter	View option	Standard		
Masking character option	X for protein, n for nucleotide	Masking col	or option	Black ▼	
☐ Show CDS translation	Align				

Sequence 1: gi|1469860|dbj|BAA09787.1|leptin [Homo sapiens] >gi|4557715|ref|NP_000221.1| leptin precursor [Homo sapiens] >gi|730218|sp|P41159|LEP_HUMAN Leptin precursor (Obesity factor) (Obese protein) >gi|623332|gb|AAA60470.1| ob >gi|1226244|gb|AAC50400.1| obese protein >gi|1407583|dbj|BAA09839.1| ob protein [Homo sapiens] >gi|38174530|gb|AAH60830.1| Leptin (obesity homolog, mouse) [Homo sapiens] >gi|46854318|gb|AAH69527.1| Leptin (obesity homolog, mouse) [Homo sapiens] >gi|46854679|gb|AAH69452.1| Leptin (obesity homolog, mouse) [Homo sapiens]... Length = 167 (1 .. 167)

Sequence 2: gi|1813869|gb|AAB41786.1|leptin precursor [Ovis aries] >gi|3041703|sp|Q28603|LEP_SHEEP Leptin (Obesity factor)
Length = 146 (1 .. 146)





NOTE:Bitscore and expect value are calculated based on the size of the nr database.



```
Score = 244 \text{ bits } (624), Expect = 1e-63
 Identities = 127/146 (86%), Positives = 136/146 (93%), Gaps = 0/146 (0%)
            {\tt VPIQKVQDDTKTLIKTIVTRINDISHTQSVSSKQKVTGLDFIPGLHPILTLSKMDQTLAV}
       22
Query
                                                                             81
            VPI+KVQDDTKTLIKTIVTRINDISHTQSVSSKQ+VTGLDFIPGLHP+L+LSKMDQTLA+
            VPIRKVQDDTKTLIKTIVTRINDISHTQSVSSKQRVTGLDFIPGLHPLLSLSKMDQTLAI
Sbjct
       1
                                                                             60
Query
       82
            YQQILTSMPSRNVIQISNDLENLRDLLHVLAFSKSCHLPWASGLETLDSLGGVLEASGYS
            YQQIL S+PSRNVIQISNDLENLRDLLH+LA SKSC LP
                                                        LE+L+SLG VLEAS YS
            {\tt YQQILASLPSRNVIQISNDLENLRDLLHLLAASKSCPLPQVRALESLESLGVVLEASLYS}
Sbjct
       61
                                                                            120
Query
       142
           TEVVALSRLQGSLQDMLWQLDLSPGC
                                         167
            TEVVALSRLQGSLQDML QLDLSPGC
Sbjct 121 TEVVALSRLQGSLQDMLRQLDLSPGC
                                         146
```

CPU time: 0.02 user secs. 0.01 sys. secs 0.03 total secs. K Lambda 0.321 0.135 0.415 Gapped Lambda K 0.267 0.0410 0.140 Matrix: BLOSUM62 Gap Penalties: Existence: 11, Extension: 1 Number of Sequences: 1 Number of Hits to DB: 241 Number of extensions: 83 Number of successful extensions: 1 Number of sequences better than 10.0: 1 Number of HSP's gapped: 1 Number of HSP's successfully gapped: 1 Length of query: 167 Length of database: 1,565,033,500 Length adjustment: 124 Effective length of query: 43 Effective length of database: 1,565,033,376 Effective search space: 67296435168 Effective search space used: 67296435168 Neighboring words threshold: 9 X1: 16 (7.4 bits) X2: 129 (49.7 bits) X3: 129 (49.7 bits) S1: 41 (21.9 bits) S2: 73 (32.7 bits)